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*DB=USPT,JPAB,EPAB,DWPI; PLUR=YES; OP=ADJ*

<u>L8</u>	(lung or pulmonary) same (cancer or tumor or tumour) and (alphaV or 'alpha v')	47	<u>L8</u>
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<u>L7</u>	L6 and (alphaV or 'alpha v')	1	<u>L7</u>
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<u>L6</u>	L5 and (pulmonary or lung) same (fibrosis or fibrotic)	62	<u>L6</u>
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<u>L5</u>	rgd and (fibrosis or fibrotic)	174	<u>L5</u>
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*DB=USPT; PLUR=YES; OP=ADJ*

<u>L4</u>	(alphav or 'alpha v') and (fibrosis or fibrotic)	16	<u>L4</u>
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*DB=USPT,EPAB,DWPI; PLUR=YES; OP=ADJ*

<u>L3</u>	(alphav or 'alpha v') same (lung or pulmonary)	10	<u>L3</u>
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<u>L2</u>	(alphav or 'alpha v') same (antibod\$) and (treat\$ or therap\$ or suppress\$ or inhibit\$ or prevent\$) same (fibrosis or fibrotic)	2	<u>L2</u>
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<u>L1</u>	(alphav or 'alpha v') same (fibrosis or fibrotic)	14	<u>L1</u>
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END OF SEARCH HISTORY

**WEST**[Generate Collection](#)[Print](#)**Search Results - Record(s) 1 through 2 of 2 returned.**

1. Document ID: KR 2002015704 A DE 19929410 A1 WO 200100660 A1 AU 200062630 A NO 200106341 A EP 1189930 A1 CZ 200104484 A3 BR 200011954 A SK 200101872 A3

L2: Entry 1 of 2

File: DWPI

Feb 28, 2002

DERWENT-ACC-NO: 2001-113366

DERWENT-WEEK: 200258

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TITLE: New octapeptide compounds as alpha v beta 6 integrin inhibitors useful for treating and diagnosing heart disease, tumors, osteoporosis, fibrosis, inflammation, infection and psoriasis

INVENTOR: DIEFENBACH, B; GROTH, U ; JONCZYK, A ; ZISCHINSKY, G

PRIORITY-DATA: 1999DE-1029410 (June 26, 1999)

## PATENT-FAMILY:

PUB-NO	PUB-DATE	LANGUAGE	PAGES	MAIN-IPC
KR 2002015704 A	February 28, 2002		000	C07K007/06
DE 19929410 A1	December 28, 2000		033	C07K007/06
WO 200100660 A1	January 4, 2001	G	000	C07K007/06
AU 200062630 A	January 31, 2001		000	C07K007/06
NO 200106341 A	February 25, 2002		000	C07K000/00
EP 1189930 A1	March 27, 2002	G	000	C07K007/06
CZ 200104484 A3	April 17, 2002		000	A61K038/04
BR 200011954 A	May 7, 2002		000	C07K007/06
SK 200101872 A3	May 9, 2002		000	C07K007/06

INT-CL (IPC): A61 K 38/04; A61 K 38/08; A61 P 7/02; C07 K 0/00; C07 K 7/06
[Full](#) | [Title](#) | [Citation](#) | [Front](#) | [Review](#) | [Classification](#) | [Date](#) | [Reference](#) | [Sequences](#) | [Attachments](#) | [Claims](#) | [KMC](#) | [Drawn Desc](#) | [Image](#)

2. Document ID: NZ 502546 A WO 9907405 A1 AU 9887743 A EP 996460 A1 BR 9814040 A CN 1267224 A CZ 200000413 A3 HU 200003547 A2 MX 2000001196 A1 KR 2001022740 A JP 2001513333 W AU 739283 B US 6316601 B1 US 2001056076 A1 US 2002004482 A1

L2: Entry 2 of 2

File: DWPI

Feb 1, 2002

DERWENT-ACC-NO: 1999-167213

DERWENT-WEEK: 200214

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TITLE: Preventing and treating acute lung injury and pulmonary fibrosis - with antagonists of integrin (v6

INVENTOR: DEAN, S; ROBERT, P ; XIAOZHU, H ; HUANG, X ; PYTELA, R ; SHEPPARD, D

PRIORITY-DATA: 1997US-055060P (August 8, 1997), 1998US-0130870 (August 7, 1998),

1999US-0365695 (August 2, 1999), 2001US-0818416 (March 27, 2001)

## PATENT-FAMILY:

PUB-NO	PUB-DATE	LANGUAGE	PAGES	MAIN-IPC
NZ 502546 A	February 1, 2002		000	A61K039/395
WO 9907405 A1	February 18, 1999	E	023	A61K038/16
AU 9887743 A	March 1, 1999		000	A61K038/16
EP 996460 A1	May 3, 2000	E	000	A61K038/16
BR 9814040 A	October 3, 2000		000	A61K038/16
CN 1267224 A	September 20, 2000		000	A61K038/16
CZ 200000413 A3	November 15, 2000		000	A61K038/16
HU 200003547 A2	February 28, 2001		000	A61K038/16
MX 2000001196 A1	October 1, 2000		000	A61K038/16
KR 2001022740 A	March 26, 2001		000	A61K038/16
JP 2001513333 W	September 4, 2001		026	C12N015/09
AU 739283 B	October 11, 2001		000	A61K038/16
US 6316601 B1	November 13, 2001		000	C07K016/28
US 2001056076 A1	December 27, 2001		000	A61K048/00
US 2002004482 A1	January 10, 2002		000	A61K038/16

INT-CL (IPC): A61 K 31/70; A61 K 31/711; A61 K 38/00; A61 K 38/16; A61 K 38/17; A61 K 39/395; A61 K 45/00; A61 K 48/00; A61 P 11/00; C07 H 21/04; C07 K 14/00; C07 K 14/435; C07 K 14/47; C07 K 16/18; C07 K 16/28; C07 K 17/00; C12 N 5/10; C12 N 5/12; C12 N 15/09; C12 P 21/08; C12 P 21/08; C12 R 1:91

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Term	Documents
ALPHAV.DWPI,EPAB,USPT.	24
ALPHAVS	0
"ALPHA V".DWPI,EPAB,USPT.	0
FIBROSIS.DWPI,EPAB,USPT.	11492
FIBROSES.DWPI,EPAB,USPT.	169
FIBROTIC.DWPI,EPAB,USPT.	2315
FIBROTICS.DWPI,EPAB,USPT.	5
ANTIBOD\$	0
ANTIBOD.DWPI,EPAB,USPT.	352
ANTIBODANTIBODA.DWPI,EPAB,USPT.	1
((ALPHAV OR 'ALPHA V') SAME (ANTIBOD\$) AND (TREAT\$ OR THERAP\$ OR SUPPRESS\$ OR INHIBITS\$ OR PREVENT\$) SAME (FIBROSIS OR FIBROTIC)).USPT,EPAB,DWPI.	2

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L3: Entry 10 of 10

File: DWPI

May 26, 1998

DERWENT-ACC-NO: 1996-171609

DERWENT-WEEK: 199831

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**TITLE:** Recombinant adenovirus vectors for gene therapy - used in the treatment of glioma, melanoma and cystic fibrosis

**Basic Abstract Text (2):**

USE - The recombinant viruses and vectors may be used in gene therapy (claimed). For example, a recombinant adenovirus having a penton base molecule recognised by alpha v beta 3 receptors can be used to treat melanoma or glioma, and a recombinant adenovirus recognised by alpha 3 beta 1 receptors and expressing the cystic fibrosis transmembrane regulator (CFTR) gene can be used to treat cystic fibrosis by delivery to the epithelial cells of the lungs. The recombinant adenoviruses may also be used to treat blood related diseases, pathogenic infections including HIV infection, and angiogenesis.

**Equivalent Abstract Text (3):**

USE - The recombinant viruses and vectors may be used in gene therapy (claimed). For example, a recombinant adenovirus having a penton base molecule recognised by alpha v beta 3 receptors can be used to treat melanoma or glioma, and a recombinant adenovirus recognised by alpha 3 beta 1 receptors and expressing the cystic fibrosis transmembrane regulator (CFTR) gene can be used to treat cystic fibrosis by delivery to the epithelial cells of the lungs. The recombinant adenoviruses may also be used to treat blood related diseases, pathogenic infections including HIV infection, and angiogenesis.

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L6: Entry 2 of 62

File: USPT

Sep 10, 2002

DOCUMENT-IDENTIFIER: US 6447753 B2

TITLE: Porous particles for deep lung delivery

Brief Summary Text (5):

Inhaled aerosols have been used for the treatment of local lung disorders including asthma and cystic fibrosis (Anderson, et al., Am. Rev. Respir. Dis., 140. 1317-1324 (1989)) and have potential for the systemic delivery of peptides and proteins as well (Patton and Platz, Advanced Drug Delivery Reviews, 8:179-196 (1992)). However, pulmonary drug delivery strategies present many difficulties for the delivery of macromolecules; these include protein denaturation during aerosolization, excessive loss of inhaled drug in the oropharyngeal cavity (typically exceeding 80%), poor control over the site of deposition, irreproducibility of therapeutic results owing to variations in breathing patterns, the quick absorption of drug potentially resulting in local toxic effects, and phagocytosis by lung macrophages.

Detailed Description Text (42):

The porous particles may include a therapeutic agent for local delivery within the lung, such as agents for the treatment of asthma, emphysema, or cystic fibrosis, or for systemic treatment. For example, genes for the treatment of diseases such as cystic fibrosis can be administered.

Other Reference Publication (30):

Anderson, P.J., et al., "Effect of Cystic Fibrosis on Inhaled Aerosol Boluses," Am. Rev. Respir. Dis., 140:1317-1324 (1989).

Other Reference Publication (42):

Barrera, D.A., et al., "Synthesis and RGD Peptide Modification of a New Biodegradable Copolymer: Poly(lactic acid-co-lysine)," J. Am. Chem. Soc., 115:11010-11011 (1993).

## CLAIMS:

9. The composition of claim 4 wherein the agent is an agent for the treatment of asthma, emphysema, or cystic fibrosis.
17. The composition of claim 12 wherein the agent is an agent for the treatment of asthma, emphysema, or cystic fibrosis.
39. The method of claim 34 wherein the agent is an agent for the treatment of asthma, emphysema, or cystic fibrosis.
49. The method of claim 44 wherein the agent is an agent for the treatment of asthma, emphysema, or cystic fibrosis.

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L6: Entry 23 of 62

File: USPT

Aug 21, 2001

DOCUMENT-IDENTIFIER: US 6277812 B1

TITLE: Methods for inhibiting TGF-.beta. activity

Detailed Description Text (2):

Increased TGF-.beta. production has been found to be an important element in a number of fibrotic diseases that are characterized by an accumulation of extracellular matrix components (Border and Ruoslahti, 1992). Besides fibronectin, collagens, and tenascin (Ignatz and Massague, 1986; Varga et al., 1987; Pearson et al., 1988), TGF-.beta. also upregulates the expression of proteoglycans (Bassols and Massague, 1988). In mesangial cells both decorin and biglycan can increase as much as 50-fold after induction by TGF-.beta. (Border et al., 1990a), whereas in fibroblasts only biglycan seems to be elevated (Romaris et al., 1992; Kahari et al., 1991). Fibromodulin has not been studied in this regard. TGF-.beta. plays a pivotal role in the pathogenesis of experimentally induced glomerulonephritis, the most critical manifestation of which is the accumulation of extracellular matrix in the glomeruli (Border et al., 1990). A recent study shows that injection of recombinant decorin into glomerulonephritic rats can suppress the matrix accumulation (Border et al., 1992). The present invention indicates that fibromodulin can be even more effective in that situation. The TGF-.beta. neutralizing activities of the decorin-type proteoglycans indicates that new types of therapeutics can be developed based on these molecules.

Detailed Description Text (13):

The invention additionally provides a method of treating a pathology caused by a TGF-.beta.-regulated activity comprising contacting the TGF-.beta. with a purified polypeptide, wherein the polypeptide comprises the TGF-.beta. binding domain of a protein and wherein the protein is characterized by a leucine-rich repeat of about 24 amino acids, whereby the pathology-causing activity is prevented or reduced. While the method is generally applicable, specific examples of pathologies which can be treated include cancer, a fibrotic disease, and glomerulonephritis. In fibrotic cancer, for example, decorin can be used to bind TGF-.beta., destroying TGF-.beta.'s growth stimulating activity on the cancer cell. Other proliferative pathologies include rheumatoid arthritis, arteriosclerosis, adult respiratory distress syndrome, cirrhosis of the liver, fibrosis of the lungs, post-myocardial infarction, cardiac fibrosis, post-angioplasty restenosis, renal interstitial fibrosis and certain dermal fibrotic conditions such as keloids and scarring.

Detailed Description Text (19):

In addition, the present invention further relates to a pharmaceutical composition containing decorin or its functional equivalent, such as biglycan or fibromodulin, and a pharmaceutically acceptable carrier useful in the above methods. Pharmaceutically acceptable carriers include, for example, hyaluronic acid, and aqueous solutions such as bicarbonate buffers, phosphate buffers, Ringer's solution and physiological saline supplemented with 5% dextrose or human serum albumin, if desired. The pharmaceutical compositions can also include other agents that promote wound healing known to those skilled in the art. Such agents can include, for example, biologically active chemicals and polypeptides, including RGD-containing polypeptides attached to a biodegradable polymer as described in PCT WO 90/06767 published on Jun. 28, 1990, and incorporated herein by reference. Such polypeptides can be attached to polymers by any means known in the art, including covalent or ionic binding, for example.

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L6: Entry 59 of 62

File: USPT

Aug 5, 1997

DOCUMENT-IDENTIFIER: US 5654270 A

TITLE: Use of fibromodulin to prevent or reduce dermal scarring

Detailed Description Text (2):

Increased TGF-.beta. production has been found to be an important element in a number of fibrotic diseases that are characterized by an accumulation of extracellular matrix components (Border and Ruoslahti, 1992). Besides fibronectin, collagens, and tenascin (Ignatz and Massague, 1986; Varga et al., 1987; Pearson et al., 1988), TGF-.beta. also upregulates the expression of proteoglycans (Bassols and Massague, 1988). In mesangial cells both decorin and biglycan can increase as much as 50-fold after induction by TGF-.beta. (Border et al., 1990a), whereas in fibroblasts only biglycan seems to be elevated (Romaris et al., 1992; Kahari et al., 1991). Fibromodulin has not been studied in this regard. TGF-.beta. plays a pivotal role in the pathogenesis of experimentally induced glomerulonephritis, the most critical manifestation of which is the accumulation of extracellular matrix in the glomeruli (Border et al., 1990). A recent study shows that injection of recombinant decorin into glomerulonephritic rats can suppress the matrix accumulation (Border et al., 1992). The present invention indicates that fibromodulin can be even more effective in that situation. The TGF-.beta. neutralizing activities of the decorin-type proteoglycans indicates that new types of therapeutics can be developed based on these molecules.

Detailed Description Text (13):

The invention additionally provides a method of treating a pathology caused by a TGF-.beta.-regulated activity comprising contacting the TGF-.beta. with a purified polypeptide, wherein the polypeptide comprises the TGF-.beta. binding domain of a protein and wherein the protein is characterized by a leucine-rich repeat of about 24 amino acids, whereby the pathology-causing activity is prevented or reduced. While the method is generally applicable, specific examples of pathologies which can be treated include cancer, a fibrotic disease, and glomerulonephritis. In fibrotic cancer, for example, decorin can be used to bind TGF-.beta., destroying TGF-.beta.'s growth stimulating activity on the cancer cell. Other proliferative pathologies include rheumatoid arthritis, arteriosclerosis, adult respiratory distress syndrome, cirrhosis of the liver, fibrosis of the lungs, post-myocardial infarction, cardiac fibrosis, post-angioplasty restenosis, renal interstitial fibrosis and certain dermal fibrotic conditions such as keloids and scarring.

Detailed Description Text (19):

In addition, the present invention further relates to a pharmaceutical composition containing decorin or its functional equivalent, such as biglycan or fibromodulin, and a pharmaceutically acceptable carrier useful in the above methods.

Pharmaceutically acceptable carriers include, for example, hyaluronic acid, and aqueous solutions such as bicarbonate buffers, phosphate buffers, Ringer's solution and physiological saline supplemented with 5% dextrose or human serum albumin, if desired. The pharmaceutical compositions can also include other agents that promote wound healing known to those skilled in the art. Such agents can include, for example, biologically active chemicals and polypeptides, including RGD-containing polypeptides attached to a biodegradable polymer as described in PCT WO 90/06767, published on Jun. 28, 1990, and incorporated herein by reference. Such polypeptides can be attached to polymers by any means known in the art, including covalent or ionic binding, for example.

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L6: Entry 62 of 62

File: DWPI

Aug 14, 2002

DERWENT-ACC-NO: 1993-182240

DERWENT-WEEK: 200261

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**TITLE:** Prevention or reduction of dermal scarring - by administration of decorin or its functional equivalents bi-glycan or fibromodulin

**Basic Abstract Text (2):**

Also claimed are: a pharmaceutical compsn. comprising decorin or its functional equivalents and a carrier, e.g. hyaluronic acid; the compsn. may further comprise an RGD-contg. polypeptide attached to a biodegradable polymer; a method of treating a pathology caused by a transforming growth factor (TGF)-beta regulated activity comprising contacting the TGF-beta with a purified polypeptide comprising a TGF-beta binding domain of a protein characterised by a leucine-rich repeat of about 24 amino acids, whereby the pathology-causing activity is prevented or reduced, the protein may be e.g. decorin, biglycan or fibromodulin.

**Basic Abstract Text (3):**

**USE/ADVANTAGE** - Decorin binds and neutralises a variety of biological functions of TGF-beta, including the induction of extracellular matrix. Thus decorin can be used to prevent or reduce dermal scarring resulting from burns, skin injuries or surgery. The polypeptides can be used for treating TGF-beta pathologies such as rheumatoid arthritis, glomerulonephritic, arteriosclerosis, adult respiratory distress syndrome, cirrhosis of the liver, fibrotic cancer, fibrosis of the lungs, post-myocardial infarction, cardiac fibrosis, post-angioplasty restenosis, renal interstitial fibrosis or dermal fibrotic conditions (claimed)

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PULMONARY.DWPI,EPAB,JPAB,USPT.	26668
PULMONARIES	0
PULMONARYS	0
CANCER.DWPI,EPAB,JPAB,USPT.	93005
CANCERS.DWPI,EPAB,JPAB,USPT.	21667
TUMOR.DWPI,EPAB,JPAB,USPT.	53934
TUMORS.DWPI,EPAB,JPAB,USPT.	33289
TUMOUR.DWPI,EPAB,JPAB,USPT.	22901
TUMOURS.DWPI,EPAB,JPAB,USPT.	12841
((LUNG OR PULMONARY) SAME (CANCER OR TUMOR OR TUMOUR) AND (ALPHAV OR 'ALPHA V')).USPT,JPAB,EPAB,DWPI.	47

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**WEST**  

L8: Entry 23 of 47

File: USPT

Nov 16, 1999

DOCUMENT-IDENTIFIER: US 5985278 A  
TITLE: Anti-.alpha.V-integrin monoclonal antibody

Brief Summary Text (7):

Studies *in vivo* also implicate .alpha.V.beta.3 in melanoma development. In the murine B16-F10 melanoma system, experimental lung metastasis could be suppressed by high levels of RGD-peptides (Hardan et al., 1993; Humphries et al., 1986), potent blockers of .alpha.v-integrin function. More recently, Felding-Habermann and colleagues have shown that .alpha.V-series integrins promote subcutaneous tumor growth of M21 human melanoma in immune-deficient mice. The M21 system is elegant, and consists of a suite of cells expressing different .alpha.V-series integrins (Kieffer et al., 1991; Felding-Habermann et al., 1992; Cheresh and Spiro, 1987). The parent, M21, expresses .alpha.V.beta.3 and .alpha.V.beta.5 (Wayner et al., 1991): it attaches to vitronectin and grows as a subcutaneous tumor. M21-L, a somatic variant of M21, has no detectable .alpha.V (Cheresh and Spiro, 1987): it cannot bind vitronectin and develops slow-growing tumors. M21-L4 is a transfectant of M21-L, stably re-expressing a full length .alpha.V-chain: it binds vitronectin and grows rapidly as a subcutaneous tumor (Felding-Habermann et al., 1992). Thus the presence of cell surface .alpha.V-integrins is directly correlated with M21 subcutaneous growth.

Drawing Description Text (3):

FIG. 1: Fluorescence Activated Cell Sorter (FACS) Analysis of the Alpha-V Group Antibodies and Controls to M21 and M21L Human Melanoma Cells

Drawing Description Text (4):

Cells were incubated with 10 g ml.<sup>-1</sup> primary antibodies, stained with fluorescently labelled secondary antibodies, counter stained with propidium iodide to allow gating of necrotic cells, and 10,000 cells per sample were analyzed. The open peak represents intensity of the second layer antibody alone. The closed peak, the intensity of the specifying primary and secondary together. Vertical axis shows cells per channel, horizontal axis shows log fluorescent intensity in that channel. M21 carries surface .alpha.v integrin, M21-L has none. The pattern of staining for the alpha-V group antibodies closely matches the LM142 (.alpha.v-specific) and LM609 (.alpha.v.beta.3-specific) stainings. Especially, they react with M21 but minimally with M21-L. Antibody 9.2.27 reacts with a surface proteoglycan. 14E2 and 21H6 recognize an otherwise undefined 200 kDa melanoma surface protein. Their staining patterns are discrete from those of the alpha-V group. Especially, they react similarly with both M21 and M21-L.

Drawing Description Text (5):

FIG. 2: The Alpha-V Group Antibodies Immunoprecipitate Similar Proteins

Drawing Description Text (19):

0.32.times.106 (.diamond-solid..box-solid.) or 1.times.10.sup.6 (.diamond..quadrature.) of M21 (.diamond-solid..diamond.) or M21L (.box-solid..quadrature.) cells were injected into the tail vein of nude mice. At the time shown, groups of animals were killed and the lungs examined for tumor nodules. All the M21 mice were killed at 42 days. For the M21-L mice, 3-6 mice were sacrificed at each point. The tumor burden after 6 weeks in the M21 mice was too high to count (>>250 per lung), so T-statistics are shown for the hypothesis that an M21-L group is from the same population as the control M21 group at 42 days at the <0.001 level (\*\*) or <0.02 (\*). Where both hi and lo injected groups have the same

significance only one is shown. The bars show the mean tumor number for the groups. Vertical axis: metastases/lung; horizontal axis: days post injection.

Detailed Description Text (3):

The alpha-V Group Monoclonal Antibodies React with Integrin .alpha.v-chain

Detailed Description Text (4):

Antibody screening by ELISA on purified .alpha.v.beta.3 and .alpha.IIb.beta.3 revealed five clones, 17E6, 20A9, 23G5, 14D9.F8 and 10G2 which reacted specifically with .alpha.v.beta.3 (Table 1). These MAbs are termed "the alpha-V group". All were IgG1 isotype. In the same ELISA assay, anti-integrin antibodies of known specificity against the .alpha.v.beta.3 complex (LM609), the .alpha.v chains (LM142), the .alpha.v.beta.5 complex (P5H9), the .alpha.IIb.beta.3 complex (CP8), the .beta.3 chains (AP3) and the .beta.1 chains (P4C10), reacted as predicted from the literature (Table 1). In ELISA on fixed cells ('CELISA'), with cells expressing .alpha.v.beta.3 and .alpha.v.beta.5 (M21), .alpha.v.beta.5 but no .alpha.v.beta.3 (UCLAP3), neither .alpha.v.beta.3 nor .alpha.v.beta.5 (M21-L), and .alpha.IIb.beta.3 (M21-L-IIb), the .alpha.V group showed a reaction pattern consistent with their recognition of the .alpha.v-integrin chain and clearly distinct from a reaction with .beta.3, .beta.5, .beta.1, or other .alpha.-chains (Table 1).

Detailed Description Text (5):

The results corroborated the ELISA data with purified receptors. MAbs with specificities for .beta.3, and GpIIb were also obtained in the screen (data not shown), and these reacted in a way clearly discrete from the alpha-v group. 17E6, 14D9.F8, 20A9 and 23G5 bound .alpha.v.beta.3 with similar apparent affinity. 50% binding was achieved at .about.10-20 ng ml.sup.-1 (.about.50-100 pM-similar to LM609). 10G2 bound similar to LM142 with about 10 times lower affinity). CP8, against .alpha.IIb.beta.3 and 14E2 (see below), showed minimal binding to .alpha.v.beta.3 at concentrations up to 100 nM.

Detailed Description Text (6):

The ability of the alpha-V group to recognize native .alpha.v-integrins was tested by FACS (FIG. 1; Table 2) and by immunoprecipitation from surface labelled cells (FIG. 2). In FACS analysis (FIG. 1), the .alpha.v-expressing line (M21) reacted strongly with 17E6, 14D9.F8, 20A9, 23G5, and with the .alpha.v-defining antibodies LM142 and LM609, moderately with 10G2, and also with the control MAbs 14E2 and 21H6 and Mab 9.2.27. By contrast, .alpha.v-deficient variant (M21-L) reacted weakly with the alpha-V group and with LM142 and LM609, but showed similar reactivity as M21 with 14E2, 21H6 and 9.2.27. M21-L has an intracellular pool of .beta.3 subunits which were detected in FACS only when the cells were permeabilized (Table 2).

Detailed Description Text (7):

In FACS analysis of M21-L4 (.alpha.v-retransfected M21-L cells (Felding-Habermann et al., 1992)), the alpha-V group gave reaction patterns as on M21. The control vector transfectants, M21-L12 and the GpIIb transfectants, M21-L-IIb (Kieffer et al., 1991), showed no reactions with the alpha-V group (Table 1). UCLAP-3 adenocarcinoma reacted with the alpha-V group, with LM142 and P5H9, but not with LM609. UCLAP-3 does not express .beta.3 (see Background). The melanoma WM793 had the same reaction pattern as M21. In immunoprecipitation screening of M21 cells, the alpha-V group gave the same immunoprecipitation patterns as LM142 (anti-.alpha.v), and LM609 (anti-.alpha.v.beta.3) (FIG. 2a). A strong broad band was seen at .about.92 kDa and a weaker band at .about.145 kDa, with weak accompanying bands at .about.100 kDa, a pattern characteristic of surface labelled .alpha.v.beta.3 and .alpha.v.beta.5 integrins (Wayner et al., 1991). When compared to the precipitation patterns on M21-L, none of the alpha-V group precipitated (data from 17E6 and 20A9 are shown), and neither did LM142 or LM609. (FIG. 2c). .beta.1-specific antibodies gave similar precipitation patterns from both cell lines. In M21-L, precipitation with anti-.beta.3 antibodies gave a band at .about.92 kDa, due to intracellular .beta.3-labelled in permeable (possibly necrotic) cells. UCLAP3 (FIG. 2d) gave no precipitate with LM609, but a .about.95 kDa/145 kDa complex was precipitated, by the alpha-V group and by LM142 (FIG. 2d). In summary, ELISA, CELISA, FACS analyses and immunoprecipitations of gave consistent reaction patterns and strongly suggested that MAbs of the alpha-V group react with extracellular domains on human .alpha.v-integrin chains.

Detailed Description Text (10):

.alpha.v-integrins can function as receptors for vitronectin, so the alpha-V group was screened for their possible effects on cell attachment to vitronectin substrates. After integrin analysis by FACS, cells were tested in attachment assays (Table 2, FIG. 3). In FACS, human melanoma and carcinoma cell lines reacted similarly with the alpha-V group. The reaction with 17E6 is summarized (FIG. 3). The initial attachment to vitronectin of cells reacting in FACS with 17E6 was strongly blocked by that antibody, but only weakly affected by the control antibody 14E2 (FIG. 3). Other members of the alpha-V group were less potent (data not shown). The vigorous attachment of murine cell B16F10 on vitronectin was not affected by 17E6 and B16F10 did not react with 17E6 in FACS. As predicted (Cheresh and Harper, 1987), B16F10 attachment to vitronectin was sensitive to micromolar concentrations of RGD-peptides, suggesting the presence of functional surface .alpha.v.beta.3 (SLG and B. Diefenbach). Thus, 17E6 and the .alpha.-V group reacted with human but not mouse .alpha.v.

Detailed Description Text (16):

The invention investigated the effect of the .alpha.v-blocking antibody 17E6 on the subcutaneous development of M21 tumors in BALB/c nu/nu mice (FIG. 7). In animal models, the development of M21 tumors in nude mice has been correlated with the cell surface expression of .alpha.v-series integrins (see Background). M21 cells were subcutaneously co-injected and endotoxin-free antibodies. 17E6 consistently (4/4 experiments) blocked the subcutaneous development of M21 tumors (FIG. 7a). No tumors (0/32) have taken in the presence of 17E6, and the animals still remain tumor free--now in excess of 6 months. Control tumor take was 75-90%. Non-blocking antibodies against the .alpha.v-chain itself and control antibodies against the melanoma cell surface showed variable and inconsistent effects on tumor development. In 14E2 treated controls, take of tumors was reduced depending on experiment 30-60%, but remaining tumors grew as the untreated controls and, like the controls, these animals had pulmonary micro-metastases revealed when the lungs were brought into tissue culture (not shown). By contrast, 17E6 treated animals had neither subcutaneous tumors nor metastases in lungs, liver, kidney, spleen, colon, stomach, nor in thoracic or abdominal body cavities when sacrificed at 6 months. The .alpha.v.beta.3-deficient line M21-L grew more slowly subcutaneously than M21, and was unaffected by 17E6. M21-L controls treated with 14E2 had a take reduced in comparison to untreated animals, similar to that seen in M21 cells (FIG. 7b).

Detailed Description Text (17):

The growth of M21 and M21-L and the effect of the antibody 17E6 were compared in an "experimental metastasis" tail-vein injection model. M21 formed many colonies in a dose dependent manner, while M21-L formed significantly fewer colonies, but did form lung nodules when injected at higher dosage (FIG. 7c). In other words, tumor growth in the lungs was also enhanced by the presence of cell surface .alpha.v-integrins, and pre-incubation of M21 with 17E6 reduced (by 90%) the numbers of tumor colonies that formed. Interestingly, the level of tumor formation was similar to those achieved by M21-L cells in the same experiment. The antibody did not alter the numbers of animals in which the tumor grew (Table 3).

Detailed Description Text (23):

Having tested the effects on cell proliferation, ADCC and AECM, it was examined whether the levels of DNA synthesis in M21 cells were affected by the MAbs. 17E6, 14E2, and LM609 at 0.5 M had no effect on thymidine incorporation (FIG. 9). DNA-synthesis in M21-L, M21-L4 and M21-L-IIb cells were also unaffected by the antibodies. M21-L and M21-L-IIb react neither with 17E6 nor LM609, but do react with 14E2 (FIG. 1). Many other melanoma cell lines were also tested and their DNA synthesis was shown not to be obviously affected by the .alpha-V group (not shown).

Detailed Description Text (39):

Tumor progression and metastasis is classically a disease where cells escape normal growth and adhesion controls and invade, migrate, attach and grow at an inappropriate site. Integrins are now known to control many cell adhesion events, and adhesion can in turn regulate mechanistically interwoven events including growth, differentiation, cell movement and the activity of protease networks, developmental events which are reiterated in the metastatic cascade (Liotta et al.,

1991; Stetler Stevenson et al., 1993; Fidler, 1988). In this study antibodies are described directed against human .alpha.V-series integrins one of which, 17E6, perturbs initial cell attachment, disrupts stable .alpha.v-ligand interactions and interferes with human melanoma development in an in vivo animal model (Fidler, 1986). In biochemical analyses the alpha-V group antibodies showed reaction patterns closely related to LM142--a well defined antibody body to human .alpha.v--but distinct from the reaction patterns of .alpha.V.beta.3-specific (LM609), .alpha.V.beta.5-specific (P5H9) and from other defined anti-integrin antibodies. Thus, the alpha-V group antibodies are likely to recognize the human .alpha.V-integrin chain. cDNA cloning of the 17E6 immunoglobulin transcripts revealed unique, unambiguous sequences. The heavy chain variable sequences were characteristic of Kabat group IIb immunoglobulins and the light chain sequences characteristic of Kabat group V (Kabat et al., 1987). Thus, the antibody is uniquely defined.

#### Detailed Description Text (43):

Growth of M21 tumors in nude mice depends strongly on .alpha.V-integrins (Felding-Habermann et al., 1992; Sanders et al., 1992). As 17E6 could modulate stable .alpha.V-ligand interactions and had a long term effect its effect on tumor development was examined. It was found that 17E6 blocked the development of subcutaneous M21 tumors in nude mice, thus strongly supporting the studies of Felding-Habermann et al. In addition, it could be shown that .alpha.v also promoted, and 17E6 inhibited, the development of M21 as experimental lung metastases. This invention has thus independently confirmed the important earlier study, and extended it by using syngeneic antibody-mediated "therapy" with 17E6. These results emphasize the importance of .alpha.V-integrins in the development of the M21 tumor, and eliminate the possibility that the earlier results arose as selection artefacts of cloning.

#### Detailed Description Text (45):

It is interesting to compare 17E6 with RGD-peptides. In the B16F10-C57blk6 murine melanoma model coinjected peptides inhibited the development of B16-F10 pulmonary tumors (Humphries et al., 1986; Hardan et al., 1993). With the same assumptions as for 17E6, .about.100 .mu.M RGD-peptide was present (Hardan et al., 1993), some two orders of magnitude over the dose required to block cell attachment to vitronectin. However, RGDS (SEQ ID NO:6) has a serum half-life of 8 min (Humphries et al., 1988). As a general therapeutic goal, it might be preferable to generate long-lived blockers for suppressing tumor development.

#### Detailed Description Text (127):

For experimental lung metastasis, cells were harvested (0.05% Trypsin/0.02% EDTA) and were injected into the tail vein of nude mice (0.5.times.10.sup.6 cells in 0.2 ml PBS++). 7 weeks later the animals were sacrificed, the lungs removed and fixed in a Bouins' solution, and the tumor foci on the surface of the lungs counted. For antibody treatment, harvested and washed cells were incubated with purified endotoxin free MAbs (70 .mu.g per 10.sup.6 cells 0.5 ml PBS++) for 30 min at 20.degree. C. in an end-over-end rotator before dilution to 0.5.times.10.sup.6 cells in 0.2 ml PBS and injection. Cells viability was assessed by Trypan blue dye exclusion before and after completing the injection schedule, where no significant difference was found (viability pre-injection=viability post injection .+- .5%). The tumor inhibition data was assessed using the 2-tailed Student T-test.

#### Detailed Description Paragraph Table (3):

TABLE 3	Inhibition of development of M21 tumor foci by 17E6 Mab in BALB/C nu/nu mice lung colonization "experimental metastasis" assay. Cells and Tumor Number of Tumor Foci % Treatment Take Mean .+- . SEM Median (Range) Control	M21 (Control) 99
87 .+- . 110 30 (3-378)	700 M21 + 17E6 78 8 .+- . 7.7 5 (0.21) 9 M21-L 56 19 .+- . 22	Table 3: M21 and M21L cells
8.5 (0-60) 22*	were harvested with trypsin/EDTA, incubated with 17E6 antibody or control antibody, washed and injected into the tail vein of nude mice. 7 weeks later the animals were sacrificed and the <u>lungs</u> examined for surface <u>tumor</u> foci. Pretreatment with 17E6 lowered the numbers of foci that developed. Similar numbers of foci developed when M21L cells (which lack .alpha.v on the cell surface) were injected. * = Compared to control: not antibodydependent.	

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**Search Results -**

<b>Term</b>	<b>Documents</b>
LUNG.DWPI,EPAB,JPAB,USPT.	46989
LUNGS.DWPI,EPAB,JPAB,USPT.	19933
PULMONARY.DWPI,EPAB,JPAB,USPT.	26668
PULMONARIES	0
PULMONARYS	0
CANCER.DWPI,EPAB,JPAB,USPT.	93005
CANCERS.DWPI,EPAB,JPAB,USPT.	21667
TUMOR.DWPI,EPAB,JPAB,USPT.	53934
TUMORS.DWPI,EPAB,JPAB,USPT.	33289
TUMOUR.DWPI,EPAB,JPAB,USPT.	22901
TUMOURS.DWPI,EPAB,JPAB,USPT.	12841
((LUNG OR PULMONARY) SAME (CANCER OR TUMOR OR TUMOUR) AND (BETA6)).USPT,JPAB,EPAB,DWPI.	2

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**Set Name Query**  
side by side

**Hit Count Set Name**  
result set

*DB=USPT,JPAB,EPAB,DWPI; PLUR=YES; OP=ADJ*

<u>L9</u>	(lung or pulmonary) same (cancer or tumor or tumour) and (beta6)	2	<u>L9</u>
<u>L8</u>	(lung or pulmonary) same (cancer or tumor or tumour) and (alphaV or 'alpha v')	47	<u>L8</u>
<u>L7</u>	L6 and (alphaV or 'alpha v')	1	<u>L7</u>
<u>L6</u>	L5 and (pulmonary or lung) same (fibrosis or fibrotic)	62	<u>L6</u>
<u>L5</u>	rgd and (fibrosis or fibrotic)	174	<u>L5</u>

*DB=USPT; PLUR=YES; OP=ADJ*

<u>L4</u>	(alphav or 'alpha v') and (fibrosis or fibrotic)	16	<u>L4</u>
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*DB=USPT,EPAB,DWPI; PLUR=YES; OP=ADJ*

<u>L3</u>	(alphav or 'alpha v') same (lung or pulmonary)	10	<u>L3</u>
<u>L2</u>	(alphav or 'alpha v') same (antibod\$) and (treat\$ or therap\$ or suppress\$ or inhibit\$ or prevent\$) same (fibrosis or fibrotic)	2	<u>L2</u>
<u>L1</u>	(alphav or 'alpha v') same (fibrosis or fibrotic)	14	<u>L1</u>

END OF SEARCH HISTORY

is in DialUnits  
? b 410  
12oct02 09:20:50 User208760 Session D2187.1  
\$0.35 0.100 DialUnits File1  
\$0.35 Estimated cost File1  
\$0.35 Estimated cost this search  
\$0.35 Estimated total session cost 0.100 DialUnits

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Set Items Description  
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HIGHLIGHT set on as ''  
HIGHLIGHT set on as ''  
? begin 5,73,155,399  
12oct02 09:20:55 User208760 Session D2187.2  
\$0.00 0.073 DialUnits File410  
\$0.00 Estimated cost File410  
\$0.01 TELNET  
\$0.01 Estimated cost this search  
\$0.36 Estimated total session cost 0.173 DialUnits

*✓✓fibrosis*

SYSTEM:OS - DIALOG OneSearch  
File 5:Biosis Previews(R) 1969-2002/Oct W1  
(c) 2002 BIOSIS  
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(c) 2002 Elsevier Science B.V.  
\*File 73: Alert feature enhanced for multiple files, duplicates removal, customized scheduling. See HELP ALERT.  
File 155:MEDLINE(R) 1966-2002/Oct W1  
\*File 155: Alert feature enhanced for multiple files, duplicates removal, customized scheduling. See HELP ALERT.  
File 399:CA SEARCH(R) 1967-2002/UD=13715  
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\*File 399: Use is subject to the terms of your user/customer agreement.  
Alert feature enhanced for multiple files, etc. See HELP ALERT.

Set Items Description  
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? s (beta6 or beta(w)6) and (fibrosis or fibrotic)  
536 BETA6  
1764082 BETA  
4130995 6  
3667 BETA(W)6  
186165 FIBROSIS  
16604 FIBROTIC  
S1 43 (BETA6 OR BETA(W)6) AND (FIBROSIS OR FIBROTIC)  
? rd s1  
...completed examining records  
S2 26 RD S1 (unique items)  
? t s2/3/all

2/3/1 (Item 1 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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13692381 BIOSIS NO.: 200200321202  
Transforming growth factor beta (TGFbeta) dependent and independent pathways of induction of tubulointerstitial fibrosis in alphavbeta6 -/- mice.

AUTHOR: Ma L-J(a); Donnert E; Sheppard D; Fogo A B  
AUTHOR ADDRESS: (a)Dept. of Pathology, Vanderbilt University, Nashville, TN  
\*\*USA  
JOURNAL: Journal of the American Society of Nephrology 12 (Program and Abstract Issue) :p819A September, 2001  
MEDIUM: print  
CONFERENCE/MEETING: ASN (American Society of Nephrology)/ISN (International Society of Nephrology) World Congress of Nephrology San Francisco, CA, USA October 10-17, 2001  
ISSN: 1046-6673  
RECORD TYPE: Citation  
LANGUAGE: English

2/3/2 (Item 2 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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13546734 BIOSIS NO.: 200200175555  
Spontaneous emphysema and upregulation of MMP-12 expression following **beta6** integrin inactivation.  
AUTHOR: Morris David G(a); Kaminski Naftali; Huang Xiaozhu; Shapiro Steven D; Sheppard Dean  
AUTHOR ADDRESS: (a)Department of Medicine, Lung Biology Center, University of California, San Francisco, San Francisco, CA, 94143\*\*USA  
JOURNAL: Molecular Biology of the Cell 11 (Supplement) :p257a Dec., 2000  
MEDIUM: print  
CONFERENCE/MEETING: 40th American Society for Cell Biology Annual Meeting San Francisco, CA, USA December 09-13, 2000  
ISSN: 1059-1524  
RECORD TYPE: Citation  
LANGUAGE: English

2/3/3 (Item 3 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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13533818 BIOSIS NO.: 200200162639  
Molecular and structural consequences of early renal allograft injury.  
AUTHOR: Baboolal Keshwar(a); Jones Geraint A; Janezic Alenka; Griffiths David R; Jurewicz Wieslaw A  
AUTHOR ADDRESS: (a)Welsh Transplant Research Group, University Hospital of Wales, Heath Park, Cardiff, CF14 4XW\*\*UK E-Mail: Baboolalk@cf.ac.uk  
JOURNAL: Kidney International 61 (2):p686-696 February, 2002  
MEDIUM: print  
ISSN: 0085-2538  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English

2/3/4 (Item 4 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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13339516 BIOSIS NO.: 200100546665  
Increased expression of tenascin-C-binding epithelial integrins in human bullous keratopathy corneas.  
AUTHOR: Ljubimov Alexander V(a); Saghizadeh Mehrnoosh; Pytela Robert; Sheppard Dean; Kenney M Cristina  
AUTHOR ADDRESS: (a)Ophthalmology Research Laboratories, Cedars-Sinai Medical Center, 8700 Beverly Boulevard, Davis-5069, Los Angeles, CA, 90048: ljubimov@cshs.org\*\*USA

JOURNAL: Journal of Histochemistry and Cytochemistry 49 (11):p1341-1350  
November, 2001  
MEDIUM: print  
ISSN: 0022-1554  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English  
SUMMARY LANGUAGE: English

2/3/5 (Item 5 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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12921523 BIOSIS NO.: 200100128672  
Protection against unilateral ureteral obstruction (UUO) induced tubulointerstitial **fibrosis** in alphavbeta6 -/- mice.  
AUTHOR: Ma L-J(a); Donnert E; Sheppard D; Fogo A B  
AUTHOR ADDRESS: (a)Department of Pathology, Vanderbilt University, Nashville, TN\*\*USA  
JOURNAL: Laboratory Investigation 81 (1):p189A January, 2001  
MEDIUM: print  
CONFERENCE/MEETING: Annual Meeting of the United States and Canadian Academy of Pathology Atlanta, Georgia, USA March 03-09, 2001  
ISSN: 0023-6837  
RECORD TYPE: Citation  
LANGUAGE: English  
SUMMARY LANGUAGE: English

2/3/6 (Item 6 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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12661639 BIOSIS NO.: 200000415141  
Mesenchymal regulation of alveolar repair in pulmonary **fibrosis**.  
AUTHOR: Fang Kenneth C(a)  
AUTHOR ADDRESS: (a)Cardiovascular Research Institute, University of California, San Francisco, San Francisco, CA, 94143-0911\*\*USA  
JOURNAL: American Journal of Respiratory Cell and Molecular Biology 23 (2):p142-145 August, 2000  
MEDIUM: print  
ISSN: 1044-1549  
DOCUMENT TYPE: Article  
RECORD TYPE: Citation  
LANGUAGE: English  
SUMMARY LANGUAGE: English

2/3/7 (Item 7 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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12410227 BIOSIS NO.: 200000163729  
Global analysis of gene expression in pulmonary **fibrosis** reveals distinct programs regulating lung inflammation and **fibrosis**.  
AUTHOR: Kaminski Naftali; Allard John D; Pittet Jean F; Zuo Fengrong; Griffiths Mark JD; Morris David; Huang Xiaozhu; Sheppard Dean; Heller Renu A(a)  
AUTHOR ADDRESS: (a)Roche Bioscience, 3401 Hillview Avenue, Palo Alto, CA, 94304\*\*USA  
JOURNAL: Proceedings of the National Academy of Sciences of the United States of America. 97 (4):p1778-1783 Feb. 15, 2000  
ISSN: 0027-8424

DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English  
SUMMARY LANGUAGE: English

2/3/8 (Item 8 from file: 5)  
DIALOG(R) File 5:Biosis Previews(R)  
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11896894 BIOSIS NO.: 199900143003  
The integrin alphavbeta6 binds and activates latent TGFbeta1: A mechanism for regulating pulmonary inflammation and fibrosis.  
AUTHOR: Munger John S; Huang Xiaozhu; Kawakatsu Hisaaki; Griffiths Mark J D ; Dalton Stephen L; Wu Jianfeng; Pittet Jean-Francois; Kaminski Naftali; Garat Chrystelle; Matthay Michael A; Rifkin Daniel B; Sheppard Dean(a)  
AUTHOR ADDRESS: (a)Lung Biology Center, Department Medicine, University California San Francisco, San Francisco, CA\*\*USA  
JOURNAL: Cell 96 (3):p319-328 Feb. 5, 1999  
ISSN: 0092-8674  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English

2/3/9 (Item 9 from file: 5)  
DIALOG(R) File 5:Biosis Previews(R)  
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11092846 BIOSIS NO.: 199799713991  
Expression of integrin cell adhesion receptors during human airway epithelial repair in vivo.  
AUTHOR: Pilewski Joseph M(a); Latoche Joseph D; Arcasoy Selim M; Albelda Steven M  
AUTHOR ADDRESS: (a)Div. Pulmonary Med., Univ. Pittsburgh, 440 Scaife Hall, 3550 Terrace St., Pittsburgh, PA 15261\*\*USA  
JOURNAL: American Journal of Physiology 273 (1 PART 1):pL256-L263 1997  
ISSN: 0002-9513  
RECORD TYPE: Abstract  
LANGUAGE: English

2/3/10 (Item 10 from file: 5)  
DIALOG(R) File 5:Biosis Previews(R)  
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10773331 BIOSIS NO.: 199799394476  
Inactivation of the **beta-6** integrin subunit gene protects against bleomycin-induced pulmonary fibrosis.  
AUTHOR: Griffiths Mark; Huang Xiao-Zhu; Wu Jian Geng; Sheppard Dean  
AUTHOR ADDRESS: Lung Biol. Cent., Univ. Calif., San Francisco, CA 94143\*\* USA  
JOURNAL: Molecular Biology of the Cell 7 (SUPPL.):p166A 1996  
CONFERENCE/MEETING: Annual Meeting of the 6th International Congress on Cell Biology and the 36th American Society for Cell Biology San Francisco, California, USA December 7-11, 1996  
ISSN: 1059-1524  
RECORD TYPE: Citation  
LANGUAGE: English

2/3/11 (Item 11 from file: 5)  
DIALOG(R) File 5:Biosis Previews(R)  
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10714773 BIOSIS NO.: 199799335918  
Induction of interstitial pneumonia in autoimmune mice by intratracheal administration of superantigen staphylococcal enterotoxin B.  
AUTHOR: Shinbori Toshifumi; Matsuki Misae; Suga Moritaka; Kakimoto Kiichi; Ando Masayuki(a)  
AUTHOR ADDRESS: (a) First Dep. Internal Med., Kumamoto Univ. Sch. Med., 1-1-1 Honjo, Kumamoto 860\*\*Japan  
JOURNAL: Cellular Immunology 174 (2):p129-137 1996  
ISSN: 0008-8749  
RECORD TYPE: Abstract  
LANGUAGE: English

2/3/12 (Item 12 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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10431140 BIOSIS NO.: 199699052285  
Cytokine gene expression in cirrhotic and non-cirrhotic human liver.  
AUTHOR: Llorente Luis(a); Richaud-Patin Yvonne; Alcocer-Castillejos Natasha ; Ruiz-Soto Rodrigo; Mercado Miguel Angel; Orozco Hector; Gamboa-Dominguez Armando; Alcocer-Varela Jorge  
AUTHOR ADDRESS: (a) Dep. Immunology Rheumatology, Instituto Nacional de la Nutricion Salvador Zubiran, Vasco de Quir\*\*Mexico  
JOURNAL: Journal of Hepatology 24 (5):p555-563 1996  
ISSN: 0168-8278  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English

2/3/13 (Item 13 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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10293223 BIOSIS NO.: 199698748141  
Interleukin 1 and tumor necrosis factors in obese alcoholics compared with normal-weight patients.  
AUTHOR: Bunout Daniel(a); Munoz Carlos; Lopez Marcelo; Pia De La Maza Maria ; Schlesinger Liana; Hirsch Sandra; Pettermann Margarita  
AUTHOR ADDRESS: (a) INTA, Univ. Chile, P.O. Box 138-11, Santiago\*\*Chile  
JOURNAL: American Journal of Clinical Nutrition 63 (3):p373-376 1996  
ISSN: 0002-9165  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English

2/3/14 (Item 14 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
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09114307 BIOSIS NO.: 199497122677  
Rapid detection of single nucleotide deletions: Application to the beta 6 (-A) mutation of the beta-globin gene and to cystic fibrosis.  
AUTHOR: Romey Marie-Catherine; Aguilar-Martinez Patricia; Demaille Jacques; Claustres Mireille(a)  
AUTHOR ADDRESS: (a) INSERM U249/CNRS UPR 9008, Laboratoire de Biochimie Genetique, Institut de Biologie, Boulevard H\*\*France  
JOURNAL: Human Genetics 92 (6):p627-628 1993  
ISSN: 0340-6717  
DOCUMENT TYPE: Article

RECORD TYPE: Abstract

LANGUAGE: English

2/3/15 (Item 15 from file: 5)  
DIALOG(R) File 5:Biosis Previews(R)  
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08802020 BIOSIS NO.: 199395091371  
Quantitation and structures of oligosaccharide chains in human trachea  
mucin glycoproteins.  
AUTHOR: Sangadala Sreedhara; Bhat U Ramadas; Mendicino Joseph(a)  
AUTHOR ADDRESS: (a)Dep. Biochem., University Georgia, Athens, Georgia 30602  
JOURNAL: Molecular and Cellular Biochemistry 118 (1):p75-90 1992  
ISSN: 0300-8177  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English

2/3/16 (Item 1 from file: 73)  
DIALOG(R) File 73:EMBASE  
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11254342 EMBASE No: 2001269685  
Integrin-mediated activation of transforming growth factor-beta<sup>1</sup> in  
**pulmonary fibrosis**  
Sheppard D.  
Dr. D. Sheppard, Lung Biology Center, UCSF Box 0854, San Francisco, CA  
94143 United States  
Chest ( CHEST ) (United States) 2001, 120/SUPPL. (49S-53S)  
CODEN: CHETB ISSN: 0012-3692  
DOCUMENT TYPE: Journal ; Conference Paper  
LANGUAGE: ENGLISH SUMMARY LANGUAGE: ENGLISH  
NUMBER OF REFERENCES: 28

2/3/17 (Item 1 from file: 155)  
DIALOG(R) File 155:MEDLINE(R)

12993001 21843125 PMID: 11854220  
Pulmonary inflammation induced by *Pseudomonas aeruginosa*  
lipopolysaccharide, phospholipase C, and exotoxin A: role of interferon  
regulatory factor 1.  
Wieland Catharina W; Siegmund Britta; Senaldi Giorgio; Vasil Michael L;  
Dinarello Charles A; Fantuzzi Giamilia  
Department of Medicine, University of Colorado Health Sciences Center,  
Denver, Colorado 80262, USA.  
Infection and immunity (United States) Mar 2002, 70 (3) p1352-8,  
ISSN 0019-9567 Journal Code: 0246127  
Contract/Grant No.: AI-15614; AI; NIAID; HL62608; HL; NHLBI  
Document type: Journal Article  
Languages: ENGLISH  
Main Citation Owner: NLM  
Record type: Completed

2/3/18 (Item 2 from file: 155)  
DIALOG(R) File 155:MEDLINE(R)

11302659 21344899 PMID: 11451914  
Integrin-mediated activation of transforming growth factor-beta(1) in  
**pulmonary fibrosis.**  
Sheppard D

Lung Biology Center, Center for Occupational and Environmental Health, Cardiovascular Research Institute, Department of Medicine, University of California, San Francisco, San Francisco, CA 94143, USA.  
Chest (United States) Jul 2001, 120 (1 Suppl) p49S-53S, ISSN 0012-3692 Journal Code: 0231335  
Document type: Journal Article  
Languages: ENGLISH  
Main Citation Owner: NLM  
Record type: Completed

2/3/19 (Item 3 from file: 155)  
DIALOG(R) File 155: MEDLINE(R)

10155488 99148265 PMID: 10025398  
The integrin alpha v beta 6 binds and activates latent TGF  
beta 1: a mechanism for regulating pulmonary inflammation and  
fibrosis.  
Munger J S; Huang X; Kawakatsu H; Griffiths M J; Dalton S L; Wu J; Pittet  
J F; Kaminski N; Garat C; Matthay M A; Rifkin D B; Sheppard D  
Department of Medicine, and Kaplan Cancer Center, New York University  
School of Medicine, New York 10016-6402, USA.  
Cell (UNITED STATES) Feb 5 1999, 96 (3) p319-28, ISSN 0092-8674  
Journal Code: 0413066  
Contract/Grant No.: HL47412; HL; NHLBI; HL53949; HL; NHLBI; HL56385; HL;  
NHLBI; +  
Document type: Journal Article  
Languages: ENGLISH  
Main Citation Owner: NLM  
Record type: Completed

2/3/20 (Item 1 from file: 399)  
DIALOG(R) File 399: CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

136084699 CA: 136(6)84699x PATENT  
Treatment of acute lung injury and fibrosis with antagonists of avbeta6  
INVENTOR(AUTHOR): Huang, Xiaozhu; Sheppard, Dean; Pytela, Robert  
LOCATION: USA  
PATENT: U.S. Pat. Appl. Publ. ; US 20020004482 A1 DATE: 20020110  
APPLICATION: US 130870 (19980807) \*US PV55060 (19970808)  
PAGES: 9 pp., Cont.-in-part of U.S. Provisional 55,060. CODEN: USXXCO  
LANGUAGE: English CLASS: 514012000; A61K-038/16A; A61K-031/70B;  
C07K-017/00B; C07K-014/00B

2/3/21 (Item 2 from file: 399)  
DIALOG(R) File 399: CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

136015216 CA: 136(2)15216a PATENT  
Methods for identifying modulators of the interaction between LAP  
(latency associated peptide) and integrin .alpha.v.beta.3, and medical use  
thereof  
INVENTOR(AUTHOR): Ludbrook, Steven; Barry, Simon; Horgan, Carmel; Miller,  
David  
LOCATION: UK,  
ASSIGNEE: Glaxo Group Limited  
PATENT: PCT International ; WO 200190186 A1 DATE: 20011129  
APPLICATION: WO 2001GB2352 (20010525) \*GB 200012991 (20000526) \*GB  
2001286 (20010105)  
PAGES: 44 pp. CODEN: PIXXD2 LANGUAGE: English CLASS: C07K-014/705A;  
C07K-014/475B; G01N-033/68B; G01N-033/566B; A61K-035/00B; A61K-038/00B

DESIGNATED COUNTRIES: AE; AG; AL; AM; AT; AU; AZ; BA; BB; BG; BR; BY; BZ; CA; CH; CN; CO; CR; CU; CZ; DE; DK; DM; DZ; EC; EE; ES; FI; GB; GD; GE; GH; GM; HR; HU; ID; IL; IN; IS; JP; KE; KG; KP; KR; KZ; LC; LK; LR; LS; LT; LU; LV; MA; MD; MG; MK; MN; MW; MZ; NO; NZ; PL; PT; RO; RU; SD; SE; SG; SI; SK; SL; TJ; TM; TR; TT; TZ; UA; UG; US; UZ; VN; YU; ZA; ZW; AM; AZ; BY; KG; KZ; MD; RU; TJ; TM DESIGNATED REGIONAL: GH; GM; KE; LS; MW; MZ; SD; SL; SZ ; TZ; UG; ZW; AT; BE; CH; CY; DE; DK; ES; FI; FR; GB; GR; IE; IT; LU; MC; NL; PT; SE; TR; BF; BJ; CF; CI; CM; GA; GN; GW; ML; MR; NE; SN; TD; TG

2/3/22 (Item 3 from file: 399)  
DIALOG(R)File 399:CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

134160854 CA: 134(12)160854v JOURNAL  
Transgenic and knockout mouse models of pulmonary inflammatory diseases  
AUTHOR(S): Griffiths, Mark  
LOCATION: Adult Intensive Care Unit, Royal Brompton Hospital, London, UK,  
SW3 6NP  
JOURNAL: Biomed. Health.Res. DATE: 2000 VOLUME: 34 NUMBER: Acute Lung  
Injury: From Inflammation to Repair PAGES: 93-104 CODEN: BIHREN ISSN:  
0929-6743 LANGUAGE: English PUBLISHER: IOS Press

2/3/23 (Item 4 from file: 399)  
DIALOG(R)File 399:CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

134086549 CA: 134(7)86549f PATENT  
Preparation of cyclic peptides for use as inhibitors of integrin  
.alpha.v.beta.6  
INVENTOR(AUTHOR): Jonczyk, Alfred; Diefenbach, Beate; Goodman, Simon  
LOCATION: Germany,  
ASSIGNEE: Merck Patent G.m.b.H.  
PATENT: Germany Offen. ; DE 19933173 A1 DATE: 20010118  
APPLICATION: DE 19933173 (19990715)  
PAGES: 20 pp. CODEN: GWXXBX LANGUAGE: German CLASS: C07K-007/64A;  
A61K-038/12B

2/3/24 (Item 5 from file: 399)  
DIALOG(R)File 399:CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

134042449 CA: 134(4)42449u PATENT  
Synthesis of peptide inhibitors of integrin .alpha.v.beta.6  
INVENTOR(AUTHOR): Jonczyk, Alfred; Diefenbach, Beate; Groth, Ulrich;  
Zischinsky, Gunther  
LOCATION: Germany,  
ASSIGNEE: Merck Patent G.m.b.H.  
PATENT: Germany Offen. ; DE 19929410 A1 DATE: 20001228  
APPLICATION: DE 19929410 (19990626)  
PAGES: 34 pp. CODEN: GWXXBX LANGUAGE: German CLASS: C07K-007/06A;  
A61K-038/08B

2/3/25 (Item 6 from file: 399)  
DIALOG(R)File 399:CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

133068988 CA: 133(6)68988y PATENT  
Integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use  
INVENTOR(AUTHOR): Diefenbach, Beate; Jonczyk, Alfred; Kraft, Sabine;  
Mehta, Ray

LOCATION: Germany,  
ASSIGNEE: Merck Patent G.m.b.H.  
PATENT: PCT International ; WO 200037487 A1 DATE: 20000629  
APPLICATION: WO 99EP9842 (19991211) \*DE 19858587 (19981219)  
PAGES: 37 pp. CODEN: PIXXD2 LANGUAGE: German CLASS: C07K-007/06A;  
C07K-007/08B; C12N-015/10B; A61K-038/04B; A61P-007/02B  
DESIGNATED COUNTRIES: AE; AL; AM; AT; AU; AZ; BA; BB; BG; BR; BY; CA; CH;  
CN; CU; CZ; DE; DK; EE; ES; FI; GB; GD; GE; GH; GM; HR; HU; ID; IL; IN; IS;  
JP; KE; KG; KP; KR; KZ; LC; LK; LR; LS; LT; LU; LV; MD; MG; MK; MN; MW; MX;  
NO; NZ; PL; PT; RO; RU; SD; SE; SG; SI; SK; SL; TJ; TM; TR; TT; UA; UG; US;  
UZ; VN; YU; ZA; ZW; AM; AZ; BY; KG; KZ; MD; RU; TJ; TM  
DESIGNATED REGIONAL: GH; GM; KE; LS; MW; SD; SL; SZ; TZ; UG; ZW; AT; BE;  
CH; CY; DE; DK; ES; FI; FR; GB; GR; IE; IT; LU; MC; NL; PT; SE; BF; BJ; CF;  
CG; CI; CM; GA; GN; GW; ML; MR; NE; SN; TD; TG

2/3/26 (Item 7 from file: 399)  
DIALOG(R) File 399:CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

130195758 CA: 130(15)195758m PATENT  
Treatment of acute lung injury and fibrosis with antagonists of  
.alpha.v.beta.6  
INVENTOR(AUTHOR): Huang, Xiaozhu; Sheppard, Dean; Pytela, Robert  
LOCATION: USA  
ASSIGNEE: The Regents of the University of California  
PATENT: PCT International ; WO 9907405 A1 DATE: 19990218  
APPLICATION: WO 98US16439 (19980807) \*US 55060 (19970808)  
PAGES: 23 pp. CODEN: PIXXD2 LANGUAGE: English CLASS: A61K-038/16A;  
A61K-038/17B; A61K-039/395B; A61K-048/00B; C07H-021/04B; C07K-014/435B;  
C07K-014/47B; C07K-016/18B; C07K-016/28B DESIGNATED COUNTRIES: AL; AM; AT;  
AU; AZ; BA; BB; BG; BR; BY; CA; CH; CN; CU; CZ; DE; DK; EE; ES; FI; GB; GE;  
GH; GM; HR; HU; ID; IL; IS; JP; KE; KG; KP; KR; KZ; LC; LK; LR; LS; LT; LU;  
LV; MD; MG; MK; MN; MW; MX; NO; NZ; PL; PT; RO; RU; SD; SE; SG; SI; SK; SL;  
TJ; TM; TR; TT; UA; UG; UZ; VN; YU; ZW; AM; AZ; BY; KG; KZ; MD; RU; TJ; TM  
DESIGNATED REGIONAL: GH; GM; KE; LS; MW; SD; SZ; UG; ZW; AT; BE; CH; CY;  
DE; DK; ES; FI; FR; GB; GR; IE; IT; LU; MC; NL; PT; SE; BF; BJ; CF; CG; CI;  
CM; GA; GN; GW; ML; MR; NE; SN; TD; TG  
?  
? t s2/7/23,24,25

2/7/23 (Item 4 from file: 399)  
DIALOG(R) File 399:CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

134086549 CA: 134(7)86549f PATENT  
Preparation of cyclic peptides for use as inhibitors of integrin  
.alpha.v.beta.6  
INVENTOR(AUTHOR): Jonczyk, Alfred; Diefenbach, Beate; Goodman, Simon  
LOCATION: Germany,  
ASSIGNEE: Merck Patent G.m.b.H.  
PATENT: Germany Offen. ; DE 19933173 A1 DATE: 20010118  
APPLICATION: DE 19933173 (19990715)  
PAGES: 20 pp. CODEN: GWXXBX LANGUAGE: German CLASS: C07K-007/64A;  
A61K-038/12B  
SECTION:  
CA234003 Amino Acids, Peptides, and Proteins  
CA201XXX Pharmacology  
CA263XXX Pharmaceuticals  
IDENTIFIERS: integrin alphav beta6 inhibitor peptide prep therapeutic  
DESCRIPTORS:  
Artery,disease...  
coronary; prepn. of cyclic peptides for use as inhibitors of integrin  
.alpha.v.beta.6 in treatment of

Peptides, preparation...  
cyclic; prepn. of cyclic peptides for use as inhibitors of integrin .alpha.v.beta.6 in treatment of disease

Heart, disease...  
infarction; prepn. of cyclic peptides for use as inhibitors of integrin .alpha.v.beta.6 in treatment of

Arteriosclerosis... Fibrosis... Infection... Inflammation... Neoplasm...  
Osteoporosis... Psoriasis... Thrombosis... Wound healing...  
prepn. of cyclic peptides for use as inhibitors of integrin .alpha.v.beta.6 in treatment of

Integrins... Receptors...  
prepn. of cyclic peptides for use as inhibitors of integrin .alpha.v.beta.6 in treatment of disease

CAS REGISTRY NUMBERS:

116821-47-7 317366-48-6P 317366-49-7P 317366-50-0P 317366-51-1P  
317366-52-2P 317366-53-3P 317366-54-4P 317366-55-5P 317366-56-6P  
317366-57-7P 317366-58-8P 317366-59-9P 317366-60-2P 317366-61-3P  
317366-62-4P 317366-63-5P 317366-64-6P 317366-65-7P 317366-66-8P  
317366-67-9P 317366-68-0P 317366-69-1P 317366-70-4P 317366-71-5P  
317366-72-6P 317366-73-7P 317366-74-8P 317366-75-9P 317366-76-0P  
317366-77-1P 317366-78-2P 317366-79-3P 317366-80-6P prepn. of  
cyclic peptides for use as inhibitors of integrin .alpha.v.beta.6 in  
treatment of

116821-47-7DP resin-bound, prepn. of cyclic peptides for use as inhibitors  
of integrin .alpha.v.beta.6 in treatment of

2/7/24 (Item 5 from file: 399)  
DIALOG(R) File 399:CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

134042449 CA: 134(4)42449u PATENT  
Synthesis of peptide inhibitors of integrin .alpha.v.beta.6  
INVENTOR(AUTHOR): Jonczyk, Alfred; Diefenbach, Beate; Groth, Ulrich;  
Zischinsky, Gunther  
LOCATION: Germany,  
ASSIGNEE: Merck Patent G.m.b.H.  
PATENT: Germany Offen. ; DE 19929410 A1 DATE: 20001228  
APPLICATION: DE 19929410 (19990626)  
PAGES: 34 pp. CODEN: GWXXBX LANGUAGE: German CLASS: C07K-007/06A;  
A61K-038/08B  
SECTION:  
CA234003 Amino Acids, Peptides, and Proteins  
CA201XXX Pharmacology  
CA263XXX Pharmaceuticals  
IDENTIFIERS: integrin alphav beta6 inhibitor peptide prepn therapeutic  
solid phase  
DESCRIPTORS:  
Artery, disease...  
coronary; prepn. of peptide inhibitors of integrin .alpha.v.beta.6 for  
treatment of disease  
Heart, disease...  
infarction; prepn. of peptide inhibitors of integrin .alpha.v.beta.6  
for treatment of disease  
Solid phase synthesis...  
peptide; prepn. of peptide inhibitors of integrin .alpha.v.beta.6 for  
treatment of disease  
Arteriosclerosis... Drug delivery systems... Drugs... Fibrosis... Infection  
... Inflammation... Integrins... Neoplasm... Osteoporosis...  
Peptides, preparation... Psoriasis... Receptors... Thrombosis... Wound  
healing...  
prepn. of peptide inhibitors of integrin .alpha.v.beta.6 for treatment  
of disease

CAS REGISTRY NUMBERS:

278777-44-9P 313245-97-5P 313246-01-4P 313246-05-8P 313246-09-2P  
313246-13-8P 313246-17-2P 313246-21-8P 313246-25-2P 313246-28-5P  
313246-31-0P 313246-34-3P 313246-39-8P 313246-41-2P 313246-44-5P  
313246-47-8P 313246-50-3P 313246-53-6P 313246-58-1P 313246-61-6P  
313246-64-9P 313246-67-2P 313246-70-7P 313246-75-2P 313246-78-5P  
313246-81-0P 313246-84-3P 313246-87-6P 313246-90-1P 313246-93-4P  
313246-96-7P 313246-99-0P 313247-02-8P 313247-05-1P 313247-09-5P  
313247-13-1P 313247-17-5P 313247-21-1P 313247-24-4P 313247-28-8P  
313247-31-3P 313247-33-5P 313247-36-8P 313247-39-1P 313247-42-6P  
313247-45-9P 313247-47-1P 313247-50-6P 313247-53-9P 313247-56-2P  
313247-59-5P 313247-62-0P 313247-65-3P 313247-68-6P 313247-72-2P  
313247-75-5P 313247-78-8P 313247-83-5P 313247-86-8P 313247-89-1P  
313247-91-5P 313247-93-7P 313247-95-9P prepn. of peptide inhibitors  
of integrin .alpha.v.beta.6 for treatment of disease  
313245-92-0DP resin-bound, prepn. of peptide inhibitors of integrin  
.alpha.v.beta.6 for treatment of disease

2/7/25 (Item 6 from file: 399)  
DIALOG(R) File 399:CA SEARCH(R)  
(c) 2002 American Chemical Society. All rts. reserv.

133068988 CA: 133(6)68988y PATENT  
Integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use  
INVENTOR(AUTHOR): Diefenbach, Beate; Jonczyk, Alfred; Kraft, Sabine;  
Mehta, Ray  
LOCATION: Germany,  
ASSIGNEE: Merck Patent G.m.b.H.  
PATENT: PCT International ; WO 200037487 A1 DATE: 20000629  
APPLICATION: WO 99EP9842 (19991211) \*DE 19858587 (19981219)  
PAGES: 37 pp. CODEN: PIXXD2 LANGUAGE: German CLASS: C07K-007/06A;  
C07K-007/08B; C12N-015/10B; A61K-038/04B; A61P-007/02B  
DESIGNATED COUNTRIES: AE; AL; AM; AT; AU; AZ; BA; BB; BG; BR; BY; CA; CH;  
CN; CU; CZ; DE; DK; EE; ES; FI; GB; GD; GE; GH; GM; HR; HU; ID; IL; IN; IS;  
JP; KE; KG; KP; KR; KZ; LC; LK; LR; LS; LT; LU; LV; MD; MG; MK; MN; MW; MX;  
NO; NZ; PL; PT; RO; RU; SD; SE; SG; SI; SK; SL; TJ; TM; TR; TT; UA; UG; US;  
UZ; VN; YU; ZA; ZW; AM; AZ; BY; KG; KZ; MD; RU; TJ; TM  
DESIGNATED REGIONAL: GH; GM; KE; LS; MW; SD; SL; SZ; TZ; UG; ZW; AT; BE;  
CH; CY; DE; DK; ES; FI; FR; GB; GR; IE; IT; LU; MC; NL; PT; SE; BF; BJ; CF;  
CG; CI; CM; GA; GN; GW; ML; MR; NE; SN; TD; TG  
SECTION:  
CA201012 Pharmacology  
CA263XXX Pharmaceuticals  
IDENTIFIERS: integrin alphav beta6 inhibitor peptide therapeutic  
DESCRIPTORS:  
Integrins...  
.alpha.v.beta.6; integrin .alpha.v.beta.6 inhibitor peptides, and  
therapeutic use  
Drug delivery systems...  
capsules; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic  
use  
Artery,disease...  
coronary; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic  
use  
Drug delivery systems...  
dragees; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic  
use  
Heart,disease...  
infarction; integrin .alpha.v.beta.6 inhibitor peptides, and  
therapeutic use  
Drug delivery systems...  
inhalants; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic  
use  
Drug delivery systems...

injections; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use

Antiarteriosclerotics... Anticoagulants... Antitumor agents...

Anti-infective agents... Anti-inflammatory agents... Cardiovascular agents ... DNA... Drug delivery systems... Envelope proteins... Fibrosis... Gene therapy... Peptides, biological studies... Psoriasis... Viral DNA... Wound healing promoters...

integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use

Drug delivery systems...

liposomes; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use

Drug delivery systems...

ointments; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use

Drug delivery systems...

solns.; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use

Drug delivery systems...

sprays; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use

Drug delivery systems...

suppositories; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use

Drug delivery systems...

tablets; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use

Osteoporosis...

therapeutic agents; integrin .alpha.v.beta.6 inhibitor peptides, and therapeutic use

CAS REGISTRY NUMBERS:

222557-93-9P 268230-24-6P 268230-25-7P 268230-26-8P 278777-32-5P  
278777-33-6P 278777-34-7P 278777-35-8P 278777-36-9P 278777-37-0P  
278777-38-1P 278777-39-2P 278777-40-5P 278777-41-6P 278777-42-7P  
278777-43-8P 278777-44-9P 278777-45-0P 278777-46-1P integrin  
.alpha.v.beta.6 inhibitor peptides, and therapeutic use

?

Set Items Description  
S1 43 (BETA6 OR BETA(W) 6) AND (FIBROSIS OR FIBROTIC)  
S2 26 RD S1 (unique items)  
? s (beta6 or beta(w)6) and (lung or pulmonary) (20n) (cancer? or tumor? or tumour?)  
Processing  
536 BETA6  
1764082 BETA  
4130995 6  
3667 BETA(W) 6  
1013660 LUNG  
770351 PULMONARY  
1910461 CANCER?  
1945134 TUMOR?  
259811 TUMOUR?  
221642 (LUNG OR PULMONARY) (20N) ((CANCER? OR TUMOR?) OR TUMOUR?)  
S3 16 (BETA6 OR BETA(W) 6) AND (LUNG OR PULMONARY) (20N) (CANCER?  
OR TUMOR? OR TUMOUR?)  
? rd s3  
...completed examining records  
S4 11 RD S3 (unique items)  
? t s4/3/all  
  
4/3/1 (Item 1 from file: 5)  
DIALOG(R) File 5:Biosis Previews(R)  
(c) 2002 BIOSIS. All rts. reserv.  
  
12862074 BIOSIS NO.: 200100069223  
A role for the integrin alphavbeta8 in the negative regulation of  
epithelial cell growth.  
AUTHOR: Cambier Stephanie; Mu De-zhi; O'Connell David; Boylen Kevin; Travis  
William; Liu Wei-hong; Broaddus V Courtney; Nishimura Stephen L(a)  
AUTHOR ADDRESS: (a)Lung Biology Center, University of California at San  
Francisco, San Francisco, CA, 94143: cdog@itsa.ucsf.edu\*\*USA  
JOURNAL: Cancer Research 60 (24):p7084-7093 December 15, 2000  
MEDIUM: print  
ISSN: 0008-5472  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English  
SUMMARY LANGUAGE: English  
  
4/3/2 (Item 2 from file: 5)  
DIALOG(R) File 5:Biosis Previews(R)  
(c) 2002 BIOSIS. All rts. reserv.  
  
10426243 BIOSIS NO.: 199699047388  
Neutrophil and cytokine activation with neonatal extracorporeal membrane  
oxygenation.  
AUTHOR: Fortenberry James D(a); Bhardwaj Vijay; Niemer Paula; Cornish J  
Devn; Wright Jean A; Bland Lee  
AUTHOR ADDRESS: (a)Dep. Pediatrics, Emory Univ. Sch. Med., 1405 Clifton Rd.  
NE, Atlanta, GA 30322\*\*USA  
JOURNAL: Journal of Pediatrics 128 (5 PART 1):p670-678 1996  
ISSN: 0022-3476  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English  
  
4/3/3 (Item 3 from file: 5)  
DIALOG(R) File 5:Biosis Previews(R)  
(c) 2002 BIOSIS. All rts. reserv.

09917682 BIOSIS NO.: 199598372600  
Expression of the **beta-6** integrin subunit in development,  
neoplasia and tissue repair suggests a role in epithelial remodeling.  
AUTHOR: Breuss J M; Gallo J; Delisser H M; Klimanskaya I V; Folkesson H G;  
Pittet J F; Nishimura S L; Aldape K; Landers D V; Carpenter W; Gillett N;  
Sheppard D; Matthay M A; Albelda S M; Kramer R H; Pytela R(a)  
AUTHOR ADDRESS: (a)Lung Biol. Cent., Dep. Med., Univ. Calif. San Francisco,  
San Francisco, CA 94143\*\*USA  
JOURNAL: Journal of Cell Science 108 (6):p2241-2251 1995  
ISSN: 0021-9533  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English

4/3/4 (Item 4 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
(c) 2002 BIOSIS. All rts. reserv.

09866238 BIOSIS NO.: 199598321156  
Distribution of integrins alpha-v-**beta-6** and alpha-9-beta-1 and  
their known ligands, fibronectin and tenascin, in human airways.  
AUTHOR: Weinacker Ann; Ferrando Ronald; Elliott Mark; Hogg James; Balmes  
John; Sheppard Dean(a)  
AUTHOR ADDRESS: (a)Lung Biol. Cent., UCSF Box 0854, San Francisco, CA 94143  
\*\*USA  
JOURNAL: American Journal of Respiratory Cell and Molecular Biology 12 (5  
) :p547-557 1995  
ISSN: 1044-1549  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English

4/3/5 (Item 5 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
(c) 2002 BIOSIS. All rts. reserv.

08762949 BIOSIS NO.: 199395052300  
T-cell subset analysis of Lewis lung carcinoma tumor rejection:  
Heterogeneity of effectors and evidence for negative regulatory  
lymphocytes correlating with metastasis.  
AUTHOR: Gelber Gohava(a); Eisenbach Lea; Feldman Michael; Goodenow Robert S  
AUTHOR ADDRESS: (a)Div. Immunol. Rheumatol., Stanford Univ. Sch. Med.,  
Stanford, Calif. 94305  
JOURNAL: Cancer Research 52 (23):p6507-6515 1992  
ISSN: 0008-5472  
DOCUMENT TYPE: Article  
RECORD TYPE: Abstract  
LANGUAGE: English

4/3/6 (Item 6 from file: 5)  
DIALOG(R)File 5:Biosis Previews(R)  
(c) 2002 BIOSIS. All rts. reserv.

04629251 BIOSIS NO.: 000079042288  
STUDIES ON SYNTHESIS AND ANTICANCER ACTIVITY OF SELECTED N-2  
FLUOROETHYL-N-NITROSOUreas  
AUTHOR: JOHNSTON T P; KUSSNER C L; CARTER R L; FRYE J L; LOMAX N R; PLOWMAN  
J; NARAYANAN V L  
AUTHOR ADDRESS: DEV. THERAPEUTICS PROGRAM, DIV. CANCER TREATMENT, NATL.  
CANCER INST., BETHESDA, MD. 20205.

JOURNAL: J MED CHEM 27 (11). 1984. 1422-1426. 1984  
FULL JOURNAL NAME: Journal of Medicinal Chemistry  
CODEN: JMCMA  
RECORD TYPE: Abstract  
LANGUAGE: ENGLISH

4/3/7 (Item 1 from file: 73)  
DIALOG(R) File 73:EMBASE  
(c) 2002 Elsevier Science B.V. All rts. reserv.

10980880 EMBASE No: 2001028793  
A role for the integrin alphavbeta3 in the negative regulation of epithelial cell growth  
Cambier S.; Mu D.-Z.; O'Connell D.; Boylen K.; Travis W.; Liu W.-H.; Courtney Broaddus C.; Nishimura S.L.  
S.L. Nishimura, Lung Biology Center, Univ. of California at San Francisco, Box 0854, San Francisco, CA 94143 United States  
AUTHOR EMAIL: cdog@itsa.ucsf.edu  
Cancer Research ( CANCER RES. ) (United States) 15 DEC 2000, 60/24 (7084-7093)  
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4/3/8 (Item 1 from file: 155)  
DIALOG(R) File 155:MEDLINE(R)

12993001 21843125 PMID: 11854220  
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Department of Medicine, University of Colorado Health Sciences Center, Denver, Colorado 80262, USA.  
Infection and immunity (United States) Mar 2002, 70 (3) p1352-8,  
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Contract/Grant No.: AI-15614; AI; NIAID; HL62608; HL; NHLBI  
Document type: Journal Article  
Languages: ENGLISH  
Main Citation Owner: NLM  
Record type: Completed

4/3/9 (Item 2 from file: 155)  
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Department of Medicine, and Kaplan Cancer Center, New York University School of Medicine, New York 10016-6402, USA.  
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Contract/Grant No.: HL47412; HL; NHLBI; HL53949; HL; NHLBI; HL56385; HL; NHLBI; +  
Document type: Journal Article  
Languages: ENGLISH  
Main Citation Owner: NLM

Record type: Completed

4/3/10 (Item 3 from file: 155)  
DIALOG(R) File 155: MEDLINE(R)

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T-cell subset analysis of Lewis lung carcinoma **tumor**  
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Gelber C; Eisenbach L; Feldman M; Goodenow R S  
Division of Immunology and Rheumatology, Stanford University School of  
Medicine, California 94305.

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Record type: Completed

4/3/11 (Item 1 from file: 399)  
DIALOG(R) File 399: CA SEARCH(R)  
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137075230 CA: 137(6)75230f PATENT

Integrin-binding domain of mitogen-activated protein kinases and its use  
for modulating cellular activity in cancer and other cells

INVENTOR(AUTHOR): Agrez, Michael Valentine

LOCATION: Australia

ASSIGNEE: The University of Newcastle Research Associates Limited

PATENT: PCT International ; WO 200251993 A1 DATE: 20020704

APPLICATION: WO 2001AU1672 (20011221) \*AU 20002305 (20001222)

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KG; KP; KR; KZ; LC; LK; LR; LS; LT; LU; LV; MA; MD; MG; MK; MN; MW; MX; MZ;  
NO; NZ; OM; PH; PL; PT; RO; RU; SD; SE; SG; SI; SK; SL; TJ; TM; TN; TR; TT;  
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DESIGNATED REGIONAL: GH; GM; KE; LS; MW; MZ; SD; SL; SZ; TZ; UG; ZM; ZW;  
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BF; BJ; CF; CG; CI; CM; GA; GN; GQ; GW; ML; MR; NE; SN; TD; TG  
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